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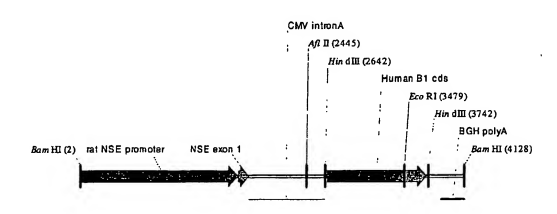
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(54) Title: TRANSGENIC RODENTS AS ANIMAL MODELS FOR MODULATION OF B₁ BRADYKININ RECEPTOR PROTEIN



NSE hB1 transgene 4132 bp

(57) Abstract: Transgenic rats are generated which incorporate a primate B₁ bradykinin receptor transgene(s) into their genome. This B₁ bradykinin receptor gene is expressed in these transgenic rats, which results in binding of compounds which are selective for the primate form (such as the human form) of the receptor and not the rat form of the receptor. Therefore, the expressed transgenes within these transgenic lines mimic antagonist and agonist selectivity of the wild type primate B₁ bradykinin receptor. These transgenic animals are useful as a specific receptor occupancy model for modulators of the B₁ bradykinin receptor from the human or closely related species, as well as providing for an animal model system for assessment of the pharmacodynamic properties of such a B₁ bradykinin modulator(s).

TITLE OF THE INVENTION TRANSGENIC RODENTS AS ANIMAL MODELS FOR MODULATION OF B₁ BRADYKININ RECEPTOR PROTEIN

5 CROSS-REFERENCE TO RELATED APPLICATIONS
Not Applicable

STATEMENT REGARDING FEDERALLY-SPONSORED R&D Not Applicable

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REFERENCE TO MICROFICHE APPENDIX Not Applicable

FIELD OF THE INVENTION

The present invention relates to transgenic rodents which express a functional B₁ bradykinin receptor protein, preferably a mammalian B₁ bradykinin receptor protein and especially a functional non-human primate or human B₁ bradykinin receptor protein. The present invention is exemplified, but in no way limited by generation of transgenic rodents wherein random integration of a DNA sequence into the rodent genome has occurred, wherein the DNA sequence encodes the open reading frame of a human B₁ bradykinin receptor protein under control of a heterologous promoter. The present invention also relates to transgenic rodents which express functional modifications of a non-human primate or human B₁ bradykinin receptor protein, including but not limited to amino acid deletions, additions, substitutions, NH2- or COOH-terminal truncations, splice variants, and the sort which provide for a protein with human B₁ bradykinin-like activity. The expressed transgenes within these transgenic lines mimic antagonist and agonist selectivity of the wild type B₁ bradykinin receptor. Therefore, the transgenic animals of the present invention are useful as a specific receptor occupancy model for modulators of a B₁ bradykinin receptor (such as a human B₁ bradykinin receptor), as well as providing for an animal model system for assessment of the pharmacodynamic properties of B1 bradykinin modulators (e.g., human B₁ bradykinin modulators), such as antagonists or agonists of receptor activity.

BACKGROUND OF THE INVENTION

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Dray and Perkins (1993, TINS 16: 99-104) and Proud and Kaplan (1988, Annual Review Immunology 6: 49-83) define two mammalian bradykinin receptor subtypes, B₁ and B₂, based on their pharmacological properties. The nonapeptide bradykinin (BK) and the decapeptide Lys-BK (kallidin) are liberated from the large protein precursor kininogen by the proteolytic action of kallikreins. BK and kallidin both activate the B₂ receptor. These B₂ receptor agonists are then degraded by a carboxypeptidase to produce the B₁ receptor agonists des-Arg⁹BK and des-Arg¹⁰kallidin or by the angiotensin converting enzyme (ACE) to yield inactive peptides. BK and kallidin act as equipotent agonists at the B₂ bradykinin receptor subtype. In contrast, BK is totally inactive at the B₁ bradykinin receptor subtype. Des-Arg10,Leu9[Kallidin] (herein, "DALK") is a peptide antagonist with structural similarities to kallidin.

The B_2 and B_1 bradykinin receptors are members of the superfamily of G-protein coupled receptors. Numerous mammalian B_1 and B_2 receptor genes have been isolated and characterized, including:

human B₁ bradykinin - U.S. Patent Nos 5,712,111 and 5,965,367, both issued to Menke et al. on January 28, 1998 and October 12, 1999, respectively, as well as Menke et al. (1994, *J. Biol. Chem.* 269:21583-21586).

rabbit B₁ bradykinin - MacNeil, et al., 1995, Biochem. Biophys. Acta 1264: 223-228.

mouse B₁ bradykinin - Hess et al., 1996, *Immunopharmacology* 33: 1-8; rat B₂ bradykinin - McEachern, et al., 1991, *Proc. Natl. Acad. Sci.* 88, 7724-7728;

human B₂ bradykinin - Hess, et al. (1992, *Biochem. Biophys. Res. Comm.* 184: 260-268); and,

rat B₁ bradykinin - Jones, et al., 1999, Eur. J. Pharmacol. 374 (3), 423-433.

Hess et al. (1996, Immunopharmacology 33: 1-8) show that B_1 receptor agonist selectivity is species specific, namely when comparing the mouse, human and rabbit B_1 receptors.

Bock and Longmore (2000, Current Opin. in Chem. Biol. 4(4):401-407) present a recent update of known modulators of B₁ and/or B₂ bradykinin receptor activity. As reviewed by the authors, it is widely held in the scientific community that B₂ receptors, but not B₁ receptors, are expressed in normal tissue. In contrast,

biologic processes which result in inflammation, pain, tissue damage can rapidly induce B_1 receptor activity, as well as bacterial infection. The apparent inducibility of the B_1 receptor under such pathological conditions may provide a therapeutic window for the use of B_1 receptor antagonists as anti-inflammatory/analgesics, thus making the B_1 receptor an attractive drug target.

To this end, there remains a need for an animal model, including but not limited to a transgenic rat model, for use as a specific receptor occupancy model for modulators of the B₁ bradykinin receptor, as well as providing for an animal model system to assess pharmacodynamic properties of potential modulators for specificity to the human B₁ bradykinin receptor. The present invention meets this ongoing need by disclosing various transgenic rodent models which express a human B₁ bradykinin receptor protein.

SUMMARY OF THE INVENTION

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The present invention relates to non-human transgenic animal cells, non-human transgenic embryos, non-human transgenic animals (including but not limited to founder animals) and/or non-human transgenic littermates, where one or more transgene(s) encoding a functional form of a non-native mammalian B_1 bradykinin receptor has been stably integrated into the germ cells and/or somatic cells of the non-human animal. Preferred non-human transgenic cells are rodent cells and a preferred non-native B_1 bradykinin receptor gene for stable integration into the rodent genome is a primate B_1 bradykinin receptor gene.

In an exemplified embodiment of the present invention, these non-human transgenic animal cells and embryos are rat cells and embryos, which subsequently give rise to a transgenic rat, including initial founder animals, littermates, and subsequent animals which comprise members of the stable transgenic line which expresses a functional form of the human B₁ bradykinin receptor. These transgenic animals contain a genetic modification such that the modified animal now expresses a functional protein which has the pharmacological properties of the human B₁ bradykinin receptor, i.e. membranes prepared from the brain of the transgenic animal (exemplified herein with transgenic rats) have pharmacological properties that are distinct from the respective non-transgenic animal.

The present invention preferably relates to animal cells wherein at least one transgene encoding a functional form of a human B₁ bradykinin receptor has been stably

integrated into the germ cells and/or somatic cells of the target animal. Additionally, the invention relates to non-human transgenic embryos, non-human transgenic founders, littermates and other transgenic animals which contain at least one transgene encoding a functional form of human B₁ bradykinin receptor. The transgenic animal cells, animals and littermates may express the non-native B₁ bradykinin receptor (e.g., a human B₁ bradykinin receptor) either in the presence or in the absence of the native (wild type) B₁ bradykinin receptor. In view of the methodology preferred for generating the transgenic rats of the present invention, a preferred transgenic cell, embryo and/or animal will contain alleles for both the native and transgenic, non-native B₁ bradykinin receptor.

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The transgenic rat models as described herein will be useful to screen any potential modulator of receptor activity (e.g., antagonists or agonists), including but not necessarily limited to peptides, proteins, or non-proteinaceous organic or inorganic molecules. To this end, the present invention relates to processes for the production of the transgenic rats of the present invention and their offspring and their use for pharmacological testing. The invention further relates to methods of determining the selectivity and activity of potential modulators of the human B₁ bradykinin receptor by administering a test compound or compounds to the transgenic rat and measuring the effect of the compound on the activity of the human B₁ bradykinin receptor. To this end, the present invention relates to various occupancy assays with, for example, brain tissue, where the ability of a test compound to penetrate the blood brain barrier, distribute into the tissue and bind to the human B₁ receptor is measured.

As used herein, the term "functional" is used to describe a gene or protein that, when present in a cell or *in vitro* system, performs normally as if in a native or unaltered condition or environment. Therefore, a gene which is not functional (i.e., "non-functional", "disrupted", "altered", or the like) will encode a protein which does not function as a wild type, native or non-altered protein, or encodes no protein at all. Such a non-functional gene may be the product of a homologous recombination event as described herein, where a non-functional gene is targeted specifically to the region of the target chromosome which contains a functional form of the gene, resulting in a "knock-out" of the wild type or native gene.

As used herein, a "modulator" is a compound that causes a change in the expression or activity of a mammalian B₂ or B₁ bradykinin receptor, such as a human B₁

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bradykinin receptor, or causes a change in the effect of the interaction of the respective receptor with its ligand(s), or other protein(s), such as an antagonist or agonist.

As used herein in reference to transgenic animals of this invention, we refer to "transgenes" and "genes". A gene is a nucleotide sequence that encodes a protein, or structural RNA. The gene and/or transgene may also include genetic regulatory elements and/or structural elements known in the art. As used and exemplified herein, a transgene is a genetic construct including a gene. The transgene is integrated into one or more chromosomes in the cells in an animal by methods known in the art. Once integrated, the transgene is carried in at least one place in the genome, preferably a chromosome, of a transgenic animal. The transgene of interest is incorporated into the target genome of the rat or other mammal, thus being introduced into their germ cells and/or somatic cells such that it is stably incorporated and is capable of carrying out a desired function. The transgene may also contain heterologous genetic regulatory elements and/or structural elements known in the art, such a heterologous promoter sequence and/or a heterologous enhancer sequence, which effects transcription of the open reading frame of the transgene within the target cell/animal. Such heterologous regulatory sequences are 'fused' or 'operatively linked' to the coding region so as to appropriately effect such gene expression. While a chromosome is the preferred target for stable incorporation of a transgene into the target animal, the term "genome" refers to the entire DNA complement of an organism, including nuclear DNA (chromosomal or extrachromosomal DNA) as well as mitochondrial DNA, which is localized within the cytoplasm of the cell. Thus, as noted previously, the transgenic rats of the present invention will stably incorporate one or more transgenes in either/or of the rat's germ cells or somatic cells (preferably both), such that the expression of the transgene (e.g., a functional form of a human B1 bradykinin gene) achieves the desired effect of presenting a specific receptor occupancy model for modulators of the human B₁ bradykinin receptor as well as providing for an pharmacodynamic animal model system to study the selectivity of test compounds to modulate the human B1 bradykinin receptor. It is preferable to introduce the transgene into a germ line cell, thereby conferring the ability to transfer the information to offspring. If offspring in fact possess some or all of the genetic information, then they, too, are transgenic animals.

As used herein, the term "animal" may include all mammals, except that when referring to transgenic animals, the use of this term excludes humans. It also includes an individual animal in all stages of development, including embryonic and fetal stages.

A "transgenic animal" is an animal containing one or more cells bearing genetic information received, directly or indirectly, by deliberate genetic manipulation at a subcellular level, such as by microinjection, targeted gene delivery such as by homologous recombination, or infection with recombinant virus. As noted above, this introduced DNA molecule (i.e., transgene) can be integrated within a chromosome, or it can be extra-chromosomally replicating DNA.

As used herein, "rodent" relates to a species which is a member of the order Rodentia, having a single pair of upper and lower incisors for gnawing, wherein the teeth grow continuously and a gap is evident between the incisors and grinding molars. Preferred examples include for generation of transgenic animals include, but are not limited to, Rattus norvegicus, Rattus rattus, and Mus musculus.

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As used herein, "rat" relates to animals which from the point of systemic zoology belong to the genus *Rattus*. The transgenic animals of the present invention may be generated from any species of the genus *Rattus*, including but not limited to *Rattus* norvegicus and *Rattus rattus*.

As used herein, "founder" refers to a transgenic animal which develops from the microinjected egg or target cell, such as an embryonic stem cell that has been targeted by a homologous recombination event to, for example, replace a rodent gene with its human homologue. The founders are tested for expression of a functional gene by any suitable assay of the gene product.

As used herein, the term "line" refers to animals that are direct descendants of one founder and bearing one transgene locus stably integrated into their germline.

As used herein, the term "inbred line" refers to animals which are genetically identical at all endogenous loci. As used in the art, inbred lines may be used for including reproducibility from one animal to the next, ability to transfer cells or tissue among animals, and the ability to carry out defined genetic studies to identify the role of endogenous genes. Such inbred lines may be developed from such lines wherein the rats that are used for microinjection are members of established inbred strains.

As used herein, the term "genotype" is the genetic constitution of an organism.

As used herein, the term "phenotype" is a collection of morphological, physiological and/or biochemical traits possessed by a cell or organism that results from the interaction of the genotype and the environment. Included in this definition of phenotype is a biochemical trait wherein a non-native transgene has been introduced into the animal, thus altering its the genotypic profile, and whereby

expression of this transgene(s) within the animal results in a new pharmacological selectivity to one or more chemical compounds, such a selectivity based on functional expression of the transgene(s) of interest. To this end, the term "phenotypic expression" relates to the expression of a transgene or transgenes which results in the production of a product, e.g., a polypeptide or protein, or alters the expression of the zygote's or the organism's natural phenotype.

As used herein, the terms "rat enolase promoter", "rat neuron specific enolase promoter", "NSE" and the such, are used interchangeably throughout this specification to refer to the promoter fragment used to exemplify the present invention, as discussed herein.

BRIEF DESCRIPTION OF THE DRAWINGS

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Figure 1 shows the transgenic plasmid targeting construct, ratEnolase intron A hB1 polyA2 [rat neuron specific enolase promoter_CMV intronA_human B1 bradykinin receptor coding sequence_BGH poly A signal CMV = human cytomegalovirus, BGH = bovine growth hormone (Construct #1; also referred to as NSE hB1)].

Figure 2A-B shows the nucleotide sequence of the integrated transgene, ratEnolase intron A hB1 polyA2, as shown pictorially in Figure 1A (SEQ ID NO:1).

Figure 3A-B shows the structure (Figure 3A) of the transcript generated from the ratEnolase intron A hB1 polyA2 targeting vector, also referred to as the NSE hB1 transcript, with the nucleotides sequence (Figure 3B) disclosed as SEQ ID NO:2.

Figure 4 shows the structure of the transgenic plasmid targeting construct, CMV promoter_CMV intron A_ human B1 cds_BGH.

Figure 5 shows a portion of the rat genomic plasmid targeting construct, CMV promoter_CMV Intron A_hB1 cds_IRES2/LacZ_BGH poly A (also referred to as pCMV B1 IZ).

Figure 6A-B shows the nucleotide sequence of the integrated transgene CMV promoter_CMV Intron A_hB1 cds_IRES2/LacZ_BGH poly A (SEQ ID NO:3).

Figure 7 shows a portion of the rat genomic plasmid targeting construct, Thy-1_hB1 cds- IZ pBS (also referred to as Thy1 hB1IZ).

Figure 8A-C shows the nucleotide sequence of the integrated transgene Thy-1_hB1 cds- IZ pBS (SEQ ID NO:4).

Figure 9A-C shows results from a saturation binding assay of ³H-DALK ([des-Arg¹⁰, Leu⁹]-Kallidin) to membranes isolated from transgenic rat brain tissue from (A): line 0004 (rat #1810); (B): line 0014 (rat #1813); and (C): line 0015 (rat #1814).

Figure 10 shows autoradiograms of brain and spinal cord sections from NSE_hB1 line 0004 transgenic rats. Non-specific binding was determined with 0.3 nM [H-3] DALK in the presence of 200nM of a non-peptide antagonist of the human B1 bradykinin receptor that has sub-nM affinity for the human B1 receptor. Total binding was determined using 0.3 nM [H-3] DALK. Regions of the brain and spinal cord that exhibit high levels of binding are indicated. Specific [H-3] DALK binding (total binding – nonspecific binding) is indicative of the level of human B1 bradykinin receptor expression. There is no detectable specific binding of [H-3] DALK in non-transgenic control rats.

15 DETAILED DESCRIPTION OF THE INVENTION

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The present invention relates to animal cells wherein at least one transgene encoding a functional form of non-native mammalian B₁ bradykinin receptor has been stably integrated into the germ cells and/or somatic cells of the target animal. To this end, the present invention relates to non-human transgenic embryos, non-human transgenic animals and non-human transgenic littermates which contain at least one transgene encoding a functional form of a non-native, mammalian B₁ bradykinin receptor. Preferred non-human transgenic cells are rodent cells and a preferred non-native B₁ bradykinin receptor gene for stable intergration into the rodent genome is a primate B₁ bradykinin receptor gene. Example of various non-human primate sources for isolated DNA molecules encoding B₁ bradykinin receptor include but are not limited to members of the old world monkey group, such as various members of the genus *Macaca*, which included *Macaca mulatta*, the rhesus monkey; members of the new world monkeys such as members of the genus *Sanuinus*, which includes the tamarins; prosimians, which include *Lemur* members, and the great apes, such as the chimpanzee (*Pan troglodytes*), orangutan (*Pongo pygmaeus*) and gorilla (*Gorilla gorilla*).

Therefore, the present invention relates to non-human transgenic animal cells, non-human transgenic embryos, non-human transgenic animals and/or non-human transgenic littermates, where one or more transgene(s) encoding a functional form of a non-native mammalian B₁ bradykinin receptor has been stably integrated into the germ

cells and/or somatic cells of the non-human animal. A preferred embodiment of this portion of the invention relates to transgenic rats which express a functional form of human B₁ bradykinin receptor, thus comprising rat transgenic cells and embryos, which subsequently give rise to a transgenic rats, including initial founder animals, littermates, and subsequent transgenic rats which represent a stable transgenic line expressing a functional form of the human B₁ bradykinin receptor. The transgene of interest contains a primate B₁ bradykinin receptor expression cassette operatively linked to regulatory sequences such as an enhancer and/or promoter fragment which are functional in the host animal, as well as a termination signal downstream of the B₁ open reading frame. The transgenic animal of the present invention allows for the investigation of pharmacological-based activity of an expressed transgene (e.g., human B₁ bradykinin receptor) in these animals, allowing for testing of the effect of certain test compounds within these transgenic animals and thus to perform preliminary tests for the development of new pharmaceutically active substances. It is evident from the data presented herein that non-human transgenic animals which incorporate a functional human B₁ bradykinin gene(s), or biologically equivalent form thereof, show a definable phenotype wherein the transgenic animal expresses an effective amount of the functional transgene product such that the transgenic animal now confers the selective pharmacological properties of the human B₁ bradykinin receptor. This phenotype is detailed herein via binding assays with membranes prepared from the brain of the transgenic rats, which are shown to have pharmacological properties that are distinct from the non-transgenic rats. This is evident in the binding data which demonstrates that human specific B₁ compounds, that have poor affinity for the rat B₁ receptor, have high affinity for membranes prepared from the brains of transgenic rats disclosed in the Example section. The endogenous rat receptor is unlikely to mask any phenotype. The radioligand ³H-DALK, has greater affinity for the human B₁ receptor than the rat B₁ receptor. This explains, in part, why no endogenous B₁ receptor activity is seen in non-transgenic rats using this ligand either by a whole brain homogenate assay or by receptor autoradiography of brain slices. In contrast, there is a binding site that is readily detectable using ³H-DALK in the exemplified transgenic rats expressing the human B₁ receptor using either of these techniques.

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The transgenic rat models as described herein will be useful to screen any potential modulator of receptor activity (e.g., antagonists or agonists), including but not necessarily limited to peptides, proteins, or non-proteinaceous organic or inorganic molecules. The transgenic animals of the present invention provide for improved models

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to study the *in vivo* effects of test compounds on human B₁ bradykinin receptor activity. Previous to this disclosure, treatments of test animal with an agent to increase wild type B₁ bradykinin expression (such as bacterial lipopolysaccharides) gave varying results to the extent in which B₁ bradykinin expression was increased and which altered the properties of the blood-brain barrier. Even if successful, the properties of compounds selective for humans could not be assessed. To circumvent this problem, attempts were made to identify a species in which the pharmacological properties of the respective species matched the human B₁ bradykinin receptor; a disadvantage being that one species may have similar properties to the human with respect to one, but not all chemical series under consideration. Alternatively, species that are closely related genetically to human (such as non-human-primates) can be used. However, this alternative suffers from the low throughput in assaying compounds through such a non-transgenic model, such as a non-human primate.

One such assay, provided only as an example and not as a limitation, is the use of the transgenic animals of the present invention in an occupancy assay in the brain to assess the ability of test compounds to penetrate the blood brain barrier as well as the ability to distribute into the tissue and bind to the receptor. A type of occupancy receptor assay may be performed using the transgenic animal of the present invention and measuring the displacement of a known radiolabeled compound which binds to the human B₁ bradykinin receptor. For example, male transgenic or non-transgenic Sprague Dawley rats may be dosed orally with test compound and fasted over night. On the day of the experiment body weights are obtained. Administration of 50%PEG/D5W vehicle (iv) or 1% methocell (po) is used to determine total binding. For iv dosed compounds, rats are placed in a perspex rat restrainer for tail vein injection of vehicle, either a test compound and/or a second compound used for determination of non-specific binding. Seven and one-half minutes later, rats are returned to the restrainer and injected with 200 µCi/kg [³H]-test compound iv. For po dosed compounds, rats are dosed by gavage with vehicle or compound. Non-specific binding is determined. Sixty minutes later, rats are placed in the restrainer and injected with 200 µCi/kg [³H]-test compound iv. Tail vein injections are through a 25 gauge, 1 inch needle on a 1 cc syringe. Tail veins are dilated by keeping rats warm and by wiping tails with an alcohol swab. Seven and 1/2 min after injection of isotope, animals are euthanized via CO₂ and the skull opened working from the base of the skull at the spinal cord opening forward to the orbital sockets. A slice of cortex

approximately 100-150 mg is cut and immediately placed in a pre-tared polypropylene tube and weighed. Cold HEPES buffer (10mM) [7.4 gms NaCl (150 mM), 4.8 gm HEPES (10 mM), 0.750 gm KCl (5 mM) per 2 liters of deionized water. pH brought to 7.4 with 1N NaOH. (NaCl & KCl: Fisher Scientific; HEPES (C₈H₁₈N₂O₄S: Boehringer Mannheim)] is added in 39 vols to each tube. Brains are 5 homogenized using a polytron homogenizer at full speed for 10 sec. Five hundred µl of homogenate are immediately filtered in duplicate through a 25mm Pall A/E filter (pre-soaked in 0.2% PEI) using a filter unit (Hoeffer). Homogenate is pipetted onto filters with valve closed so homogenate covers entire surface area of filter, valve is 10 then opened to allow filtering and washing), followed immediately by 5 x 5 ml washes of cold HEPES (5mM KCl, 150 mM NaCl, 10 mM HEPES) buffer. Each filter is placed in a scintillation vial and Ultima Gold scintillation fluid (10 ml) is added to each vial. Duplicate 500 µl aliquots of homogenate are pipetted into scintillation vials, 10 ml of Ultima Gold is added, and counted to measure total brain labelling. Samples are allowed to sit for 4 hours before counting with a tritium 15 counting program. Isotope solution is counted with the tritium program to determine actual mCi concentration. A 0.001ml sample is pipetted into a vial containing 10 ml of scintillation fluid (pipet tip is carefully wiped with a kimwipe). Calculations are as follows: (1) Percent accumulation of label (% Acc) for each sample = (Filter dpm/Homogenate dpm)x100; (2) Percent specific accumulation (% Sp Acc) for the 20 entire assay = Mean % Acc for Total Binding - Mean % Acc for Non-specific binding; (3) Percent inhibition of binding (% Inh) = ((Mean % Acc for Total Binding - % Acc for the sample))/ % Sp Acc)x100; (4) $1\mu \text{Ci} = 2.2 \text{ X } 10^6 \text{ DPM } (0.001 \text{ml})$ isotope solution(0.2mCi/ml) = 4.4 X 10⁵ DPM). Dose of the test compound may be 200 µCi/kg BW for dosing volume of 0.15 ml in 150 gm rat (0.2 mCi/ml): 1 ml of 25 1mC/ml (NEN) + 4 ml saline.

A preferred embodiment of the present invention is the generation of transgenic rodents in which one transgene encoding a functional form of a human B₁ bradykinin receptor has been stably integrated into the germ cells and/or somatic cells of the target animal. However, the present invention relates to the generation of other transgenic, non-human animals, other than the preferred targets of rats and mice, which exhibit substantially similar phenotypic traits as the exemplified transgenic rats disclosed herein, including but not limited to cows, pigs, rabbits, guinea pigs, sheep, hamsters, and goats.

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Also, the present invention preferably relates to animal cells wherein at least one transgene encoding a functional form of a human B₁ bradykinin receptor has been stably integrated into the germ cells and/or somatic cells of the target animal. Additionally, the invention relates to non-human transgenic embryos, non-human transgenic animals and non-human transgenic littermates which contain at least one transgene encoding a functional form of human B₁ bradykinin receptor. The transgenic animal cells, animals and littermates may express the non-native B₁ bradykinin receptor (e.g., a human B₁ bradykinin receptor) either in the presence or in the absence of the native (wild type) B₁ bradykinin receptor. In view of the methodology preferred for generating the transgenic rats of the present invention, a preferred transgenic cell, embryo and/or animal will contain alleles for both the native and transgenic, non-native B₁ bradykinin receptor. Typically, the transgene of interest contains a human B₁ bradykinin receptor expression cassette linked to regulatory sequences such as an enhancer and/or promoter fragment which are functional in the host animal, as well as a termination signal downstream of the B₁ open reading frame. In the preferred mode of generating a transgenic rat encoding human B₁ bradykinin receptor, the transgene is typically integrated into a host chromosomal location by nonhomologous integration. These transgenes may further comprise a selectable marker, such as a neo or gpt gene operably linked to a constitutive promoter, such as a phosphoglycerate kinase (pgk) promoter or HSV tk gene promoter linked to an enhancer (e.g., SV40 enhancer).

In some embodiments, such as the targeted insertion of the human B1 bradykinin gene into mice, the endogenous nonhuman B1 alleles are functionally disrupted so that expression of endogenously encoded murine B1 is suppressed or eliminated, so as to not interfere with expression of the human B1 transgene.

Transgenes may be incorporated into embryonic, fetal or adult pluripotent stem cells (Capecchi, 1991, Science 244: 1288-1292, see also U.S. Patent Nos. No. 5,464,764; 5,487,992; 5,627,059; 5,631,153 and 6,204,061 issued March 20, 2001) hereby incorporated by reference. Embryonic stem cells can be isolated from blastocysts cultivated *in vitro* and stably cultured within differentiation. For example, a transgene may be contained within a gene targeting vector, wherein the vector contains homologous arms (see Cappecchi, *supra*) which can be used to direct a transgene to a specific genomic site within the target ES cell. Such foreign DNA can be incorporated into the embryonic stem cells by electroporation. Embryonic stem cells which carry the transgene in the appropriate fashion are injected into the inner cell

mass of blastocysts. A chimeric animal is generated which is then crossbred to obtain animals wherein all cell carry the transgene. Along with microinjection described below, ES cell-based techniques are a preferable methodology for generating transgenic mice. A common scheme to disrupt gene function by gene targeting in ES cells is to generate a targeting construct which is designed to undergo a homologous recombination with its chromosomal counterpart in the ES cell genome. The targeting constructs are typically arranged so that they insert additional sequences, such as a positive selection marker, into coding elements of the target gene, thereby functionally disrupting it. To this end, the present invention also relates to methods of producing nonhuman animals (e.g., non-primate mammals) that have the endogenous B₁ gene inactivated by gene targeting with a homologous recombination targeting construct. General principles regarding the construction of targeting constructs and selection methods are reviewed in Bradley et al., 1992, *Bio/Technology* 10: 534, which is hereby incorporated herein by reference.

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It is within the scope of the present invention to present a transgene encoding a mammalian form of interest to the host target cell as a DNA construction such that expression of the respective B₁ bradykinin receptor is controlled by various homologous or heterologous regulatory sequences operatively linked to the B₁ bradykinin gene. To present as examples, but certainly not as a limitation, see Figure 1 (rat neuron specific enolase [NSE] promoter and CMV Intron A fused (i.e., operatively linked) to the human B₁ bradykinin receptor gene, which is upstream of the bovine growth hormone (BGH) transcriptional termination and polyadenylation signal). Animals which integrate this construct will present neuron specific expression within the central nervous system. In contrast, peripheral expression of the human B₁ bradykinin gene will occur via the integration construct shown in Figure 4 (CMV promoter/Intron A fused to the human B₁ bradykinin receptor gene, which is upstream of a second open reading frame (LacZ) which is separated by an internal ribosome entry site (IRES), with a BGH termination signal downstream of the LacZ ORF). Also, the transgene construct shown in Figure 7 (Thy-1 promoter fused to the human B₁ bradykinin receptor gene, which is upstream of a second open reading frame (LacZ) which is in turn separated by an internal ribosome entry site (IRES), with a BGH termination signal downstream of the LacZ ORF) should promote brain specific expression. These various constructions show that any of a myriad of promoter/transgene constructions may be generated for transfer into target cells for

stable, genomic DNA integration and subsequent manipulation to produce a requisite transgenic animal. The exemplified constructs described herein also provide for the integration of discistronic constructs into a non-human transgenic animal. A preferred discistronic construct utilizes an internal ribosome entry site (IRES) to separate the respective open reading frames (ORF). It is preferred that a first ORF encode for a functional form of a primate B₁ bradykinin receptor while a second ORF encode a reporter gene which allows for easy detection of tissue and/or cellular specific expression. Various reporter genes are well known in the art and include, as an example but certainly not a limitation, LacZ, green fluorescent protein (GFP), chloramphenical acetyl transferase (CAT), alkaline phosphatase and luciferase.

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A preferred method of generating a transgenic rat generally comprises first introducing DNA which includes the selected transgene into germ cells of the rat (typically fertilized eggs). These fertilized germ cells are then used to generate a complete, transgenic animal. The DNA is preferably introduced into the germ cells by known microinjection techniques, which comprises introducing the DNA into a germ cell through the aid of a microscope and a microinjector pipette which deposits intact DNA into one of the two pronuclei. Transgenic animals are selected which have incorporated into their genome at least one, and possibly more than one, selected transgene(s). At least one founder transgenic rat is selected for breeding so as to establish at least one transgenic rat line which contains the stably integrated transgene. This methodology is disclosed in U.S. Patent No. 4,873,191. Other known techniques available in the art may be utilized to generate the transgenic animals of the present invention, including but not limited to in vitro fertilization using sperm as a carrier of exogenous DNA, electroporation or alternatively, transfection into a rat embryonic stem cell line may be utilized to directly target the transgene into the rat genome, followed by selection and introduction of selected, recombinant ES cells into a rat blastocyst.. Various methodolodgy is reviewed in Mullins, et al., in Transgenic Animal: Generation and Use, Ch. 2.-Transgenic Rats, pp.7-9, Harwood Academic Publisher, 1997. Therefore, a preferred and well known method for preparing transgenic rats of the present invention includes the following steps: subjecting a female to hormonal conditions to promote superovulation (with a continuous infusion of a follicle stimulating hormone), fertilization of the superovulated female (preferably by either breeding with a fertile male or via artificial insemination), introduction of the transgene into the fertilized eggs by known techniques, such as

microinjection; implantation of the fertilized eggs into a pseudopregnant female rat, who is then brought to term. Once the fetuses in the pseudopregnant female have been brought to term, a founder animal is identified by standard techniques of hybridization of transgene DNA to genomic DNA from weanling offspring or by a PCR assay that is specific for the presence of the transgene. Founders that express the gene, particularly those that express the gene at levels and with the intended tissue distribution (such as brain specific expression) are selected and bred to establish the intended line or lines of transgenic rats.

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It will be appreciated upon practicing the present invention that not all transgenic animals which have an incorporated human B1 bradykinin gene will exhibit appropriate expression of the B₁ genes of interest. For instance, data presented in Examples 3 and 6 show variable binding of ³H-DALK to the human B₁ bradykinin receptor on three separate transgenic rat lines expressing the human B₁ bradykinin receptor. Identifying an appropriate transgenic line may also be construct specific, such a differences in promoter strength, number of transgenes incorporated into the genome, as well as the location of these integration events. The rat B₁ receptor is normally expressed at a much lower level than the transgene but its expression can be induced by certain treatments, e.g. lipopolysaccharide or streptozocin. As shown herein when comparing transgenic to non-transgenic rats, the rat B₁ bradykinin receptor has pharmacological properties that are distinct relative to the human receptor, i.e. many synthetic compounds that have high affinity for the human B₁ receptor have low affinity for the rat B₁ receptor. Animals which express the transgene at sufficient amounts under normal conditions are especially useful in receptor occupancy assays. Animals which have expression levels similar to or greater than line 0004, as measured in whole tissue assays, are preferred. However, lines with lower tissue expression (such as lines 0014 and 0015) may be useful if, for example, expression is localized within a discrete region of the tissue which is amenable to further study. To this end, one of ordinary skill in the art can expect to generate from about 6 to about 10 or so lines to be ensured that at least one of the resultant lines will exhibit the desired trait. It may also be useful to identify and breed animals which have multiple copies of the human B1 bradykinin incorporated into the target genome, such as from 2 to about 50 copies of the selected transgene. Therefore, it is within the purview of the present invention to characterize a specific transgenic animal to find a best fit for in vivolex vivo assays to determine binding

and/or receptor occupancy characteristics of for a specific test compound, wherein a specific binding/pharmacological profile will exist for the test compound in regard to the native and transgenic B₁ bradykinin receptor protein.

The nomenclature used herein and the laboratory procedures in transgenic protocols, cell culture, molecular genetics, and molecular biology are well known and commonly employed in the art. Standard techniques are used for recombinant nucleic acid methods, polynucleotide synthesis, cell culture, and transgene incorporation (e.g., electroporation, microinjection, lipofection). Various enzymatic reactions, oligonucleotide synthesis, and purification steps are performed according to the manufacturer's specifications. The techniques and procedures are generally performed according to conventional methods in the art and various general references which are provided throughout this document. The procedures therein are believed to be well known in the art and are provided for the convenience of the reader. All the information contained therein is incorporated herein by reference. The following examples are presented by the way of illustration and, because various other embodiments will be apparent to those in the art, the following is not to be construed as a limitation on the scope of the invention.

EXAMPLE 1

20 Construction of Transgenic Targeting Vectors

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Construct #1 - ratEnolase intronA hB1 polyA2 vector -

[Step 1] - Rat genomic DNA (50 or 100 ng/50 ul reaction) was used as a template to generate a PCR fragment comprising the rat neuron specific enolase promoter region. Thirty two cycles of PCR were performed (94°C 25 sec, 60°C 25 sec, 68°C 3 min) with Expand High fidelity polymerase (Roche). The forward primer was: Rat_enl.2f: 5'-CATCACTGAGCCCAACACACAA-3' (SEQ ID NO:5) and the reverse primer was Rat_enl.2r: 5'-TCACCTCGAGGACTGCAGAC-3' (SEQ ID NO:6). This PCR product was 2059 bp in length.

[Step 2] - The purified PCR product (Qiaquick PCR purification) from Step 1 was used as a template for a second round of PCR to add a BamHI restriction site. Thirty cycles of PCR (94°C 25 sec, 60°C 25 sec, 68°C 3 min) with Expand High fidelity polymerase using the following primers: Forward Rat_enl.4f, 5'-GCGGATCCTGAGCTCCTCTCTCTCTCTCGC-3' (SEQ ID NO:7); Reverse

NSE_1R, 5'-CTCGAGGACTGCAGACTCAG-3' (SEQ ID NO:8). The resulting product is 1814 bp in length.

[Step 3a] - A plasmid DNA template containing the CMVIntron A sequence was used as a template to generate a PCR fragment for subcloning. Twenty five cycles (94°C 25 sec, 60°C 25 sec, 72°C 1 min) were performed using Pfu polymerase (Stratagene). The forward primer was CMVintA.1F: 5'-GTAAGTACCGCCTATAGAGTC-3' (SEQ ID NO:9) and the reverse primer is CMVintA.1R: 5'CTGCAGAAAAGACCCATGGAAAGG-3' (SEQ ID NO:10). This PCR product is 827 bp in length.

[Step 3b] - Twelve cycles of PCR were used (94°C 25 sec, 60°C 25 sec, 68°C 1 min 10 sec with either Pfu (Stratagene) or Expand High fidelity polymerase) to add overlap ends to the CMV intron A product of Step 3a. The forward and reverse primers are as follows: Forward:NSE_CMV.OLF1:

5'-GAGTCTGCAGTCCTCGAGGTAAGTACCGCCTATAGAGTC-3' (SEQ ID NO:11);

Reverse CMV_hB1.OLR1:

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5'-TGGCGGCGGTACCAAGCTTCTGCAGAAAAGACCCATGGAAAG-3' (SEQ ID NO:12). This PCR product is 863 bp in length.

[Step 4] - A PCR fragment comprising the human B₁ bradykinin receptor coding sequence plus bovine growth hormone (BGH) polyA signal with overlap ends was constructed via 25 cycles of PCR (94°C 25 sec, 60°C 25 sec, 72°C 3 min) from plasmid pcDNA3 which contains the human bradykinin B1 receptor sequence fused to the BGH poly A sequence. The primers were as follows:

Forward CMV_hB1.OLF1:

25 5'-CTTTCCATGGGTCTTTTCTGCAGAAGCTTGGTACCGCCGCCA-3' (SEQ ID NO:13);

Reverse: BGH.1RNot:

5'- GCGCGGCCGCTCCCAGCATGCCTGCTATTG-3' (SEQ ID NO:14). This PCR is 1518 bp in length.

[Step 5] -The CMV intron A was combined with the with human B1_BGH polyA fragments via 25 cycles of PCR (94°C 25 sec, 60°C 25 sec, 68°C 4 min 30 sec) using the templates purified from Step 3 and Step 4. The primers were as follows: Forward NSE_CMV.OLF1:

5'-GAGTCTGCAGTCCTCGAGGTAAGTACCGCCTATAGAGTC-3' (SEQ ID NO:15);

Reverse: BGH.1RNot:

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5'- GCGCGGCCGCTCCCCAGCATGCCTGCTATTG-3' (SEQ ID NO:14). The PCR product of Step 5 is 2342 bp.

[Step 6] - The PCR product from step 2 was digested with Bam HI and the PCR product from Step 5 was digested with Not I. A three-way ligation was performed with BamHI/NotI digested pCR®-Blunt II-TOPO® vector (Invitrogen).

DNA sequence analysis was performed to select clones containing the fewest PCR errors. Selected clones were subcloned into BamHI/NotI digested pBlueScript (pBS) by 3-way ligation with the Bam HI/Afl II 2443 bp fragment and the Afl II/Not I 1699 bp fragment. The resulting transgene is shown in Figure 1 while the nucleotide sequence of transgene is shown in Figure 2A-B. A schematic of the transcript for this construct is shown in Figure 3A while the nucleotide sequence of the projected transcript (shown as a DNA sequence) of the transcript is shown in Figure 3B.

Construct #2 - CMV promoter_CMV intron A_human B1 cds_BGH poly A signal vector -

[Step 1- CMV promoter] One hundred nanograms of pcDNA3 was subjected to 18 cycles of PCR (94°C 25 sec, 60°C 25 sec, 72°C 1 min) with either Pfu or Expand High Fidelity polymerse. The primers were as follows:

Forward CMV promoter 1F:

5'-CGGCGCCGCCGATGTACGGGCCAGATATAC-3' (SEQ ID NO:16); Reverse:

5'-GACTCTATAGGCGGTACTTACCTATAGTGAGTCGTATTAATTTCG-3' (SEQ ID NO:17). The resulting product is 702 bp.

[Step 2 - CMV intron A] - One hundred nanograms of a DNA plasmid template comprising the CMV Intron A fragment was subjected to 18 cycles of PCR (94°C 25 sec, 60°C 25 sec, 72°C 1 min) with Pfu polymerase. The primers were as follows:

30 Forward CMV promoter_intron A 1F 5'-CGAAATTAATACGACTCACTATAGGTAAGTACCGCCTATAGAGTC-3'
(SEQ ID NO:18);
Reverse CMVintA.1R -

5'-CTGCAGAAAAGACCCATGGAAAGG-3' (SEQ ID NO:10). The product is 850 bp in length.

[Step 3] The CMV promoter fragment was linked to the CMV intronA by subjecting the PCR products of Step 1 and Step 2 to 18 cycles of PCR (94°C 25 sec,

5 60°C 25 sec, 72°C 1 min 30 sec) with Pfu polymerase. The primers were as follows:

Template: 2 ng of each PCR product from step 1 and step 2

Primers: Forward CMV promoter 1F

5'-CGGCGGCCGCCGATGTACGGGCCAGATATAC-3' (SEQ ID NO:16);

Reverse CMVintA.1R

10 5'-CTGCAGAAAAGACCCATGGAAAGG-3' (SEQ ID NO:10). This PCR product is 1508 bp in length.

[Step 4] - The CMV promoter_CMV intron A_human B1 bradykinin receptor coding sequence_ BGH poly A signal was constructed by digesting the PCR product from Step 3 with Afl II (cuts in CMV intron A). The ratEnolase intronA hB1 polyA2 vector described in this Example was digested with EcoRV and Afl II and these digested fragments were ligated together to generate the transgene shown in Figure 4.

Construct #3 - CMV intron A:human B1 coding:IRES element:Lac Z: BGH poly A - The targeting vector as detailed in Figure 5 was generated as follows:

[Step 1] - The pIRES puro plasmid (Clontech) was used as a template to
20 generate a PCR fragment comprising the IRES element. The PCR reaction was
carried out for 20 cycles (94°C 25 sec, 60°C 25 sec, 68°C 1 min 30 sec) with Expand
High Fidelity polymerase.

Primers: forward HB1_IRES F-

5'-CCAACTTTTCTGGCGGAATTAATGCATCTAGGGCGGCCAATTC-3' (SEQ

25 ID NO: 19);

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Reverse: IS_LACZ_1R -

5'-GTAAAACGACGGGATCTATCATGGTGGCGGCGGTTGGCAAGCTTA TCATCGTG-3' (SEQ ID NO:20). The resulting product is 639 bp in length. [Step 2]: The LacZ coding region was generated as a PCR fragment by utilizing

pcDNA3 beta-Gal plasmid DNA (Invitrogen) as template and running a PCR reaction for 28 cycles (94°C 25 sec, 60°C 25 sec, 68°C 3 min) with Pfu and Expand High Fidelity polymerase. The primers were as follows:

Primers: forward: IS_LACZ_1F -

5'-CACGATGATAAGCTTGCCAACCGCCGCCACCATGATAGATCCCGTC

GTTTTAC-3'(SEQ ID NO:21);

Reverse: 5'-GCCTCGAGCTATTTTTGACACCAGACCAACTG-3' (SEQ ID

NO:22). The resulting product is 3098 bp in length.

[Step 3]: The PCR products of Step 1 and Step 2 were linked via 18 cycles of

5 PCR (94°C 25 sec, 60°C 25 sec, 68°C 4 min) with Expand High Fidelity polymerase.

Primers: forward HB1_IRES F

5'-CCAACTTTTCTGGCGGAATTAATGCATCTAGGGCGGCCAATTC-3' (SEQ

ID NO: 19)

Reverse: LZ_BGH R

10 5'-CATTTAGGTGACACTATAGAATCTATTTTTGACACCAGACCAACTG-3' (SEQ ID NO:23). The resulting product is 3721 bp in length.

[Step 4] - The BGH poly A signal is generated by PCR from the plasmid pcDNA3 (Invitrogen) via 18 cycles (94°C 25 sec, 60°C 25 sec, 68°C 4 min) with Expand High Fidelity polymerase. The primers are as follows:

15 Forward LZ_BGH F -

5'-CAGTTGGTCTGGTGTCAAAAATAGATTCTATAGTGTCACCTAAATG-3' (SEQ ID NO:24);

Reverse: BGH.1RNot -

5'- GCGCGGCCGCTCCCCAGCATGCCTGCTATTG-3' (SEQ ID NO:14).

[Step 5] -The BGH polyA PCR fragment of Step 4 was linked to the IRES:LacZ fragment of Step 3 via 20 cycles of PCR (94°C 25 sec, 60°C 25 sec, 68°C 4 min 30 sec) with Expand High Fidelity polymerase. The primers are as follows: HB1_IRES F -

5'-CCAACTTTTCTGGCGGAATTAATGCATCTAGGGCGGCCAATTC-3' (SEQ

25 ID NO:19);

Reverse: BGH.1RNot -

5'- GCGCGGCCGCTCCCCAGCATGCCTGCTATTG-3' (SEQ ID NO:14)

Product: 3963 bp

[Step 6] - The ratEnolase intronA hB1 polyA2 vector described in this

30 Example was used as a template to generate a CMV intron A:human B1 coding sequence via 18 cycles (94°C 25 sec, 60°C 25 sec, 68°C 2 min 30 sec) of PCR, using the Pfu polymerase. The primers were as follows:

Forward CMVintA.1F - 5'- GTAAGTACCGCCTATAGAGTC-3' (SEQ ID NO:9);

Reverse: HB1_IRES R

5'-GAATTGGCCGCCCTAGATGCATTAATTCCGCCAGAAAAGTTGG-3' (SEQ ID NO:25). The resulting product is 1931 bp in length.

[Step 7] - The PCR products from Step 6 (CMVintron A: human B1 cds) and Step 1 (IRES) are used as a template to link these to DNA fragments by PCR.

5 Twenty cycles (94°C 25 sec, 60°C 25 sec, 68°C 4 min 30 sec) and Expand High Fidelity polymerase were utilized with the following primers:

Forward: CMVintA.1F

5'-GTAAGTACCGCCTATAGAGTC-3'(SEQ ID NO:9);

Reverse: IS_LACZ_1R

10 5'-GTAAAACGACGGGATCTATCATGGTGGCGGCGGTTGGCAAGCTTA TCATCGTG-3' (SEQ ID NO:20). The resulting product is 2549 bp in length.

[Step 8]: The PCR products from Step 7 and Step 5 are used to link the CMVintron A: human B cds (Step 7) to IRES_LacZ_BGH poly A (Step 6) via 18 cycles (94°C 25 sec, 60°C 25 sec, 68°C 7 min 30 sec) of PCR. The primers

15 were as follows:

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Forward: CMVintA.1F

5'-GTAAGTACCGCCTATAGAGTC-3 (SEQ ID NO:9)

Reverse: BGH.1RNot

5'- GCGCGGCCGCTCCCCAGCATGCCTGCTATTG-3' (SEQ ID NO:14). The resulting product is 5851 bp in length. This fragment is then subcloned into the DNA expression plasmid pCRII Topo Blunt (Invitrogen) and subjected to DNA sequence analysis to confirm generation of the appropriate transgene.

Construct 4 - CMV promoter: CMV intron A:human B1 coding: IRES2 element: Lac Z: BGH poly A -

[Step 1] A 520 bp Bgl II/Nsi I fragment from Construct 3 is subcloned into pIRES2-EGFP (Clontech). This subclone is digested with Bgl II/Nco I. A PCR fragment spanning a portion of LacZ is generated from a pcDNA3_beta Gal (Invitrogen) template via 37 cycles of PCR (94°C 25 sec, 60°C 25 sec, 68°C 2 min 30 sec) with Pfu polymerase. The primers were:

30 Forward: LZ_BspHI -

5'-GGCATCATGATAGATCCCGTCGTTTTAC-3' (SEQ ID NO:26);

Reverse: 5'- IZ_2699R

5'-TACTGTGAGCCAGAGTTGCC-3' (SEQ ID NO:27). The resulting product is 2081 bp in length. This product is digested with BspHI and EcoRV. This 1113 bp

fragment and the Bgl II/Nco I fragment above are ligated with Construct #2 and digested with Bgl II and EcoRV. The target transgene is shown schematically in Figure 5 and the nucleotide sequence of the transgene is shown in Figure 6A-B.

Construct #5 - Thy-1 promoter:human B1 coding:IRES2 element:Lac Z: BGH 5 poly A -

[Step 1] - A DNA fragment comprising the mouse Thy-1 promoter was generated from a PCR reaction using mouse genomic DNA as a template. The PCR reaction was carried out for 30 cycles (94°C 25 sec, 60°C 25 sec, 68°C 3 min 30 sec) with Expand High Fidelity polymerase. The primers were as follows:

10 Forward - Thy1_1f_Not:

5'-GCGCGGCCGCTCTGGTTATCCAGGCTTCTG-3' (SEQ ID NO:28);

Reverse - Thy1_hB1r:

5'-GGTGGCGGCGGTACCAAGCTTGTGCCAAGAGTTCCGACTTG-3' (SEQ ID NO:29). The resulting PCR product is 2923 in length.

[Step 2] - A portion of human B₁ Bradykinin coding sequence was generated from 10 ng human B₁ receptor cloned into pcDNA3. PCR conditions were as follows: 18 cycles of PCR (94°C 25 sec, 60°C 25 sec, 68°C 4 min) with Pfu polymerase. The primers were:

Forward - Thy1_hB1f:

20 5'-CAAGTCGGAACTCTTGGCACAAGCTTGGTACCGCCGCCACC-3' (SEQ ID NO:30);

Reverse: hB1_2r

5'- TGCTTGCACCTGGAATAAG-3' (SEQ ID NO:31). The resulting product was 881 bp in length.

[Step 3] -The PCR products from Step 1 and Step 2 were linked via a PCR reaction (18 cycles @ 94°C 25 sec, 60°C 25 sec, 68°C 4 min) with Expand High Fidelity polymerase. The primers were:

Forward - Thy1_1f_Not:

5'-GCGCGGCCGCTCTGGTTATCCAGGCTTCTG-3' (SEQ ID NO:28)

30 Reverse - hB1_2r:

5'- TGCTTGCACCTGGAATAAG-3' (SEQ ID NO:32). The resulting product is 3753 bp in length and was cloned into the TOPO TA vector (Invitrogen), followed by DNA sequence analysis of clones.

[Step 4] - The clone from Step 3 is digested with Not I/Bgl II and the 3473 bp fragment is isolated. Construct #4 is digested with Bgl II/Not I and the 4451 bp fragment is isolated. These two fragments are ligated into Not I digested pBlueScript (pBS), resulting the in the transgene disclosed schematically in Figure 7 and via the nucleotide sequence in Figure 8A-C.

EXAMPLE 2

Generation of Transgenic Rats Expressing Human B₁ Bradykinin 1 Receptor

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Approximately 20 ug of NSE promoter_CMV intronA_human B1 (Figure 1) cloned into pBluescript was digested with Bam HI. The 4.1 kb insert was separated from the 3 kb vector on a 0.8 % agarose gel. The 4.1 kb band was excised and extracted using Qiaquick Gel Extraction (Qiagen), following extraction the fragment was further purified by separation on a 0.8 % agarose gel. The band was excised and extracted from the gel as before with the modification of twice purifying on the Quiquick columns. The final product was resuspended in 10 mM Tris pH 7.4, 0.1 mM EDTA at a concentration of approximately 50 ng/ul. The CMV (Figure 4) and the Thy-1 promoter constructs (Figure 7) were prepared in a similar manner with the exception that Not I digestion was used to excise the linear DNA fragment for microinjection from the vector.

A purified NSE promoter_CMV intronA_human B1 (Construct #1, Figure 1A) fragment was transferred to DNX Transgenic Sciences (Now Xenogen Corporation) in Princeton, NJ under contract for the generation of transgenic rats containing this transgene. Standard methodology is utilized to transfer said construct into Sprague-Dawley rat eggs to create transgenic rat lines (see, e.g., U.S. Patent No. 4,873,191) which have incorporated at least one copy of the transgene into the genome. Three such transgenic lines which were transferred and subjected to further genomic and phenotypic analysis were lines 0004, 0014 and 0015. Line 0004 is estimated to have approximately 10 copies, with 0014 having more than line 0004. Of course, while there is a relationship, copy number and expression level are in general poorly correlated.

A Taqman assay was developed for the transcript resulting from transgenic insert containing the NSE promoter_CMV intronA_human B1 bradykinin receptor coding sequence_BGH poly A signal. The splicing of CMV intronA results in a transcript which includes 118 nucleotides of exon 1 of the neuron specific enolase

gene fused to the human B1 bradykinin receptor coding sequence (Figure 3A). PCR primers were designed such that the 3' end of the forward primers, either NSE_TM1f 5'-GAGTCTGCAGTCCTCGAGAAGC-3' (SEQ ID NO:33) or NSE_TM2f 5'-TGAGTCTGCAGTCCTCGAGAAG-3' (SEQ ID NO:34), corresponded to the spliced transcript and therefore would not detect either unspliced transcript or genomic DNA. Taqman probes, NSE_TAQ1, 5'-CTCCAATCCTCCAACCAGAGCCAGC-3' (SEQ ID NO:35), and NSE_TAQ2, 5'-TCCAATCCTCCAACCAGAGCCAGCT-3' (SEQ ID NO:36)labeled with FAM and TAMRA were designed to detect the PCR products.

An Oligotex Direct mRNA kit (Qiagen) was used to prepare mRNA from the brain of 2 transgenic and 1 non-transgenic rat from line 004. Products derived from the transgenic construct were detected using an ABI PRISM 7700 Sequence Detection System with rodent GAPDH utilized as an internal control. Rodent GAPDH was detected in all samples in contrast the product derived from the transgene was only detected in the transgenic animals. This indicates that the transcript derived from the transgenic insert in line 004 is correctly processed and that this assay can be utilized to distinguish transgenic from non-transgenic animals.

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Rat genomic DNA was prepared from tissue by proteinase K digestion followed by phenol chloroform extraction and ethanol precipitation. The genomic DNA (5 to 10 ug) was digested with Eco RI and fragments separated on a 1 % agarose gel. DNA was transferred from the gel to Zeta-Probe Genomic Tested blotting membranes (BioRad) using a VacuGene system (Pharmacia Biotech). Pfu polymerase was used to amplify a 701 nucleotide PCR product from the transgenic construct with the forward primer CMV_381F 5'- AATCTCGGGTACGTGTTCCG-3' (SEQ ID NO:37) and reverse primer Enl_gt2r 5'- TTGGCCAGGTAGATTTCTGC-3' (SEQ ID NO:38). The product was purified by Qiaquick PCR purification (Qiagen) and radiolabeled with alpha³²PdCTP by random prime labeling (Roche). Hybridization was performed in 0.25 M Na₂HPO₄, 6.5% SDS, and 10% dextran sulfate at 65°C overnight. The blot washed with a final wash of 0.1X SSC 0.1% SDS for 30 minutes at 60°C and exposed to film. There is a single Eco RI site in the NSE promoter construct therefore digestion yields a unit length band of 4132 nucleotides, similarly the CMV promoter construct of 6522 contains a single Eco RI site.

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EXAMPLE 3

Ligand Binding to Human B₁ Bradykinin Receptor Purified from Transgenic Rats Three of the five lines (0004, 0014, and 0015) of transgenic rats containing Construct #1 (neuron-specific enolase promoter driving expression of the human B₁ bradykinin receptor), as described in Example Section 2, were tested for the ability to bind ³H-DALK, a compound which is approximately 40 fold selective for this ligand. Also important in these assays is the low expression level of endogenous B₁ receptor in neuronal tissue. Briefly, transgenic animals from line 0004, 0014, and 0015 (all females) were decapitated following anesthesia and the whole brain was removed, bisected sagitally and the entire ½ brain weighed. Weights were as follows: line 0004 (813 mg), line 0014 (851 mg), and 0015 (843 mg). The brain tissue was homogenized with a Polytron in ice cold 50mM Tris·HCl, 1mM EDTA, 1mM o-phenanthroline, pH 7.4. The homogenate was centrifuged at 50,000 x g for 20 minutes. The pellet was resuspended and homogenized a second time in Tris buffer, and the centrifugation step was repeated. The final pellet was resuspended in assay buffer (20 mM HEPES, 120 mM NaCl, 5 mM KCl, 1 mM o-phenanthroline, 0.2uM of enaliprilat (the diacid form and active metabolite of enalipril which is added to inhibit angiotensin converting enzyme), 100 μg/ml bacitracin, 3 μM amastatin, 1 μM phosphoramidon, 0.1% BSA, pH 7.4. The assay was carried out in a 0.5 ml volume at room temperature for sixty minutes with 10 mg wet weight tissue/tube. Total protein was determined using a Bio Rad DC assay kit. Specific binding is measured as that which is sensitive to competition with a B₁ specific ligand, either cold DALK or a compound with specificity for the human B₁ receptor. Figure 9A, 9B and 9C show measurements of the amount of total, nonspecific and specific binding of ³H-DALK to transgenic rat brain tissue which expresses human B₁ bradykinin receptor. Line 0004 (Figure 9A) shows expression of 40 fmol/mg protein, Line 0014 (Figure 9B) shows expression of 4 fmol/mg protein, while Line 0015 (Figure 9C) shows expression of 7 fmol/mg protein. In contrast, no B₁ receptor is detected in the brains of non-transgenic rats. The Ki values determined for three standard lead compounds in Line 0004 are very similar to those obtained at the cloned hB1 receptor expressed in CHO cells. Therefore, expression of the human B₁ bradykinin receptor in Line 0004 has the properties expected for the human B₁ receptor.

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Line 0004 was subjected to autoradiographic study of the expression of human bradykinin B₁ receptor in transgenic rat brain and spinal cord. A transgenic rat (line 0004) was first anesthetized, and then the brain was removed and immediately frozen on dry ice. The coronal sections (20µm) of the brain were prepared in a cryostat. The adjacent sections of selected brain regions were divided into two sets and pre-incubated for 15 minutes at room temperature (RT) in buffer A. Following pre-incubation, two sets of the brain sections were incubated separately in buffer B for 90 min at RT. One set of the sections was incubated with 0.3 nM of [3H]DALK, and another set was incubated with both 0.3 nM of [3H]DALK and 200 µM of unlabeled L-864747. At the end of the incubation period, sections were washed three times (4 min each) in ice-cold buffer A and then rinsed in ice-cold deionized water for five seconds. Sections were dried by cold air at room temperature, then placed in a cassette against Fuji Imaging Plate (BAS-TR2025) at room temperature for a week. The plate was scanned with Fuji BAS-5000 machine, and the images were analyzed using the MCID M5 software (Imaging Research Inc.). Buffer A is 50 mM Tris-HCl, pH 7.5, 120 mM NaCl, 5 mM KCl and Buffer B is 50 mM Tris-HCl, pH 7.5, 120 mM NaCl, 5 mM KCl, 100 μg/ml Bacitracin, Sigma B-0125, 1 μM Phosphoramidon, Sigma R-7385, 1 mM o-Phenanthroline, Sigma P-9375, 3 µM Amastatin, Sigma A-1276, 0.1% BSA (Sigma A-7030). [3H]DALK is purchased from NEN Life Science (Cat.# NET1096).

The purpose of this autoradiographic study is to characterize human bradykinin B₁ receptor expression in the spinal cord and brain tissues of the transgenic rat carrying human bradykinin B₁ receptor gene by autoradiography. The radiotracer, [³H]DALK for the B₁ receptor was employed in the study and an antagonist of human bradykinin B₁ receptor was used to block the receptor specific binding of [³H]DALK. A signal that was not competed by the antagonist was defined as nonspecific binding of [³H]DALK. The results of autoradiographic study demonstrate expression of human bradykinin B₁ receptor in the brain and spinal cord of the transgenic rat. In NSE_human B₁ receptor transgenic line 0004, the expression of human bradykinin B₁ receptor varies among the different regions of the brain and spinal cord examined. The highest binding signals for [³H]DALK in transgenic rats are in the dorsal horn of the spinal cord, the cerebral cortex, hypothalamus, thalamus, cerebellum, substantial nigra, interpeduncular nucleus, nucleus of solitary tract, periaqueductal gray, and pontine nucleus. In contrast, [³H]DALK did not show any

specific binding signal in the corresponding regions of the brain and spinal cord of the non-transgenic rats, showing that integration of the human B₁ bradykinin gene into the rat genome confers a phenotype of non-native, selective binding characteristics to various test compounds and known modulators of the human B₁ bradykinin receptor.

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EXAMPLE 4

Mapping of the transgene integration site for NSE-hB1 line 0004

Genomic DNA was prepared from tissue of a transgenic rat from line 0004. The genomic DNA was partially digested with restriction endonuclease Sau 3A1 and 10 cloned into the superCOS I vector according to the manufacturer's instructions (Stratagene, La Jolla CA). Cosmid clones were screened by standard in situ hybridization of bacterial colonies using a radiolabeled probe consisting of 701 nucleotides. The probe was obtained using standard PCR conditions with the primers, 5'-AATCTCGGGTACGTGTTCCG 3' (SEQ ID NO:39) and 5' -TTGGCCAGGTAGATTTCTGC 3' (SEQ ID NO:40), and the NSE-hB1 transgene 15 construct as the template. Positive colonies were re-screened and cosmid DNA was prepared from clones that were positive in the secondary screen. Cosmid end sequencing was performed using T3 and T7 primers. DNA sequence of cosmid clone 19 that was obtained with the T3 primer was found to match rat genomic DNA containing a portion of the pellucidae glycoprotein gene 1 (ZP-1), whereas the 20 sequence from the T7 primer matched a portion of the NSE_hB1 transgene construct. To fine map the transgene integration site, cosmid 19 was digested with the restriction endonuclease DraI, and the resulting fragments were sub-cloned into the vector pBluescript II (Stratatene, La Jolla, CA). Plasmid DNA was prepared from ampicillin resistant colonies and the size of the insert was determined, clones with various size 25 inserts were analyzed by DNA sequence analysis using m13 forward and reverse primers. DNA sequence analysis of clone Dra37 revealed that it contained rat genomic DNA and a portion of the NSE-hB₁ transgene construct. Thus clone Dra37 contained one end of the transgene insertion site. BioInformatic analysis of rat genomic DNA sequence from Dra37 indicated that it matched the DNA sequence of 30 Rattus norvegicus clone CH230-6B11 (GenBank Accession number AC097387). The clone CH230-6B11 contains the zona pellucidae glycoprotein gene 1 (ZP1), the same gene that was identified by end sequencing of cosmid clone 19, and is mapped to chromosome 1. Therefore the transgene integrated into chromosome 1 near the ZP1

gene. The delineation of the transgene insertion site permitted the development of a genotyping assay for identification of the line 0004 homozygous transgenic rats. This randon integration site contains mutliple copies of the transgene. The sequence of clone Dra 37 containing the line 0004 transgene insertion site is as follows: GTAGCCTGCC TCCGATATTT GTTAGAACAA CGGTTCCCCG CCACCTACCA ACTGTTTATG TTTTCTCTAA CAAAAACCAG ACCGGCCGCT GGGCCTGATA CCTGAGTTCA GTCACCAAGA CCCACGTGGC AGAAGGAGA AACTGACTTC TGCATATTAT CCTCCAACAC ACACACAC TGGTGGCACA TTGAGAACTT ACCTCAGAAA AAAGGTAAGT AGATAAAGTA AAACTAAAAT 10 GGAGTGAGTC ACACTGGAGT TCCATGTTAC CAAATTAAAA CTAGCTTTCT GACCTTCTGA GAAACCAGGA CAGAAAGAGG TGAAGGCCAC ATTTTCTAGC CATGCCAACT GCAGCAAACA TAACTCTGTT CTGGCTGCCA TTGTCCTTAT GAAAAGTAAG CAGGAGGGAT CTGATCTATT AACCAGCTAG CTCTGTGCTT CCCTCCTCTT CTCCCAACCT CCCAAGGAAA ACATACTCCG TCCTTTTCCT TTGTTTTATT CCTGCTTCCT GTCTAGGAAA TCACTCCCCT CCAAGGCGTC AGAACACATT CTGGCTTACA GAATGAAGTT TTACCCAATT CTAGAATCAC AAAATATAGC 15 CAACGTAAAC CTTGAATGTG ATCTAATTGG TCTAAGAGGC AGAAATGAGA TGAAGAAAAA AACTGCCGAC ATAGATITCA GTCTATGGGA TGATGGGCAC ATAAACAATA AGAAGAAAGT GCCAGACAGG GGTAGGTGCT CTAAATACAA GATAAATTAG AGCAGGTTGA GAAGATGGTA CTGGGGATTG GAGGGCGAC TGCTTTAGGC AGGGTATGGG AAAGGTATGC CCCCTGAGAG 20 AGGATGTTCA TTTTTAGCAC TTGAATTTTA TTTTAGTGTA TGTGTATGCA TGTGCCACAG CAAATGTATA GAAGTAAAAG GAGACCTTGA GAGAAGTGGT TCACTCCTCC CATGTTGGTC TTGGGATCGA AGTCAGGTTG TTAGACTTGA CAGGAAGTTT CTCTCCCCAG TGAGCTGTCT CACCAGCCCA AAGGGTGGCA ACATTTTTGC TGAGACCTAA ATAAAGGACA TGCGTCAGTT CAGAAACCAC AGATATCTGA TCAACCAAGC TCCTGCAGTC TCACCTCATC TTCCTCTCAG 25 CCACACTGGC CCTTCAGTGG CCCCAGCAGT CCCCGAGGTA GGTGGCTCAA AATGTTTATG TGGCTACCTT TCATCAACTC CTTCCCCATC TCCAGCCCCG GCCAGACCCT CCAGGGCAAA CTGAGGCCTC ATCTGAGCTC CTCCTCTGCT CGCCCAATCC TTCCAACCCC CTATGGTGGT ATTGTCTGTT TACCCTATAG GACATCCTAT AGGGTAAACA GACAATAGAC CATAGGACAA CAGGCAGGAG CATGCCTGCT ATTGTCCTCC CTTGTCCTCC CTGCCATCCT AAAGCTGGCA 30 GGTGGCTGGT GGTATATGGA GGATGTAGCT GGGCCAGGGA AAAGATCCTG CACTAAAAAT CTGAAGCTAA AAATAACAGG ACACGGGATG GAGGAGCTCA GGTGGTATGG CTGACACAGA AAATGTCTGC TCCTGTATGG GACATTTGCC CCTCTTCTCC AAATATAAGA CAGGATGAGG CCTAGCTTTT GCTGCTCCAA AGTTTTA (SEQ ID NO:41).

EXAMPLE 5

Development of a Genotype Assay for NSE_hB₁ line 0004 Homozygous Transgenic Rats

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The genomic DNA sequence upstream of the transgene insertion site was utilized to design forward, 5'-GAGGTGAAGGCCACATTTTCTAGC -3' (SEQ ID NO:42), and reverse 5'- ATGGGGAAGGAGTTGATGAAAGGTAGCC -3' (SEQ ID NO:43), PCR primers. Using the cosmid DNA template and standard PCR procedures these primers generate a product of 922 nucleotides. This fragment of 922 nucleotides serves an external probe that can be radiolabeled and used in Southern blot analysis to discern wild type from transgenic chromosomes. Accordingly, Southern blot analysis of wild type rat genomic DNA with the restriction endonuclease DraI results in the detection of a single fragment of approximately 3.1 kb with the external probe. In contrast, the digestion of genomic DNA prepared from a rat heterozygous for the transgene results in the detection of two fragments, one of 3.1kb and a second of approximately 1.6kb. The 1.6 kb fragment corresponds to the chromosome with the transgene insertion site, whereas the 3.1 kb fragment corresponds to the wild type chromosome. Thus, DraI digestion and Southern blot analysis with the external probe can be used to identify homozygous wild type animals, heterozygous and homozygous transgenics. This was used to identify and establish a homozygous transgenic breeding colony. Significantly, the line 0004 homozygous animals express 2-fold more human B₁ bradykinin receptor in the brain and spinal cord than the heterozygous animals.

EXAMPLE 6

25 Ex vivo Receptor Occupancy Assay in NSE hB₁ transgenic rat

Transgenic rats of either sex are placed in an induction chamber and anesthetized with isoflurane under a Flow Sciences hood. Once anesthetized, the rat is placed on a circulating water warming blanket (Gaymar T-pump) and anesthesia is maintained using 2% isoflurane by means of a nose cone. The tail vein is cannulated with a 25G winged infusion set-up connected to a syringe containing either test compound or vehicle. The desired dose of test compound is administered. At the experimental end-point a blood sample is taken, the rat is euthanized, and tissue is removed (typically brain and spinal cord) for subsequent assays.

For autoradiographic analysis of human B₁ receptor expression, tissues removed from transgenic rats were frozen on dry ice powder, and stored at - 70°C. Coronal sections of the brain and the transverse sections of the spinal cord were prepared with cryostat (Leica, CM3050) at 20 µM of each. The frozen sections were stored at -70°C. For analysis, frozen sections were warmed at room temperature (RT) for 15 minutes, then followed by 15 minutes preincubation in the buffer without radioligand at RT. After preincubation, the sections were transferred to the incubation buffer, and incubated for 90 minutes at RT. Total binding, both non-specific and specific, was determined by incubating in buffer containing 0.3 nM [H-3] DALK. An adjacent section was utilized to determine non-specific binding, which was incubated in buffer containing 0.3 nM [H-3] DALK and 200 nM of a non-peptide receptor antagonist that exhibits high affinity and specificity for the human B₁ bradykinin receptor. Following the 90 minute incubation, the sections were washed three times, 3 minutes each, in buffer, rinsed in DIH₂O for 30 seconds at 4°C, and then dried by air blower at RT. The sections were placed against Fuji imaging plates, and exposed for a week at RT. The plates were scanned with Fuji PhosphorImager BAS 5000, and the images were analyzed with MCID M5 software. Figure 10 shows autoradiograms of brain and spinal cord sections from NSE_hB1 line 0004 transgenic rats. Regions of the brain and spinal cord that exhibit high levels of binding are indicated. Specific [H-3] DALK binding (total binding – nonspecific binding) is indicative of the level of human B₁ bradykinin receptor expression. There is no detectable specific binding of [H-3] DALK in non-transgenic control rats.

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For homogenate-based binding assay, thirty-five milligrams of frozen brain (cerebral cortex or cerebellum) or spinal cord is homogenized with a Polytron, in a large volume of ice-cold assay buffer (20mM HEPES, 120mM NaCl, 5mM KCl, pH 7.4) and transferred to two chilled centrifuge tubes. To pellet membranes the tubes are centrifuged for 10 minutes at 75,000xg in a rotor pre-cooled to 4°C. The supernatant is discarded and each tube is rinsed with 20ml ice-cold buffer and then homogenized pellets above in ice-cold assay buffer. The homogenate is pooled and added to a tube containing the radiotracer, 20pM of a non-peptide human B₁ receptor antagonist that is labeled with ³⁵S, in each tube containing 0.5ml room temperature assay buffer. Nonspecific binding is determined by adding homogenate to tubes containing the radiotracer and 100nM of the unlabeled non-peptide human B₁ receptor antagonist. At set time points (1,2,4,6,8,10 minutes) the contents of three tubes are

filtered over individual 25mm GF/B filters presoaked in 0.05% Triton X-100. The filtration step is performed by adding 4ml ice-cold assay buffer to each of the three replicate tubes, pouring the contents over the filters, and washing each filter two times with 4ml ice-cold buffer. A Hoeffer FH 225V filtration manifold is used for the filtration. The nonspecific binding tubes are similarly filtered after finishing the 6 time points. Filters are transferred to 5ml scintillation vials and counted after soaking 10 hours in 3ml Beckman Ready Safe scintillation fluid.

The specific binding is calculated at each time point (total cpm – nonspecific cpm) and the slope of the association is determined by linear regression. Receptor occupancy in a drug treated animal is determined by the following equation:

% Occupancy = $(1-(slope_{drug}/slope_{vehicle})) \times 100$ slope_{drug} is the slope of the association rate line from a drug treated animal. slope_{vehicle} is the slope determined for a vehicle treated animal.

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Various patent and journal publications are cited herein, the disclosures of which are all incorporated by reference in their entireties.

WHAT IS CLAIMED IS:

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A non-human transgenic animal having incorporated into its genome at least one copy of a transgene encoding a primate bradykinin B₁ receptor gene, or a
 functional equivalent thereof, such that the transgenic animal demonstrates a humanized B₁ bradykinin receptor occupancy or binding profile.

- 2. A non-human transgenic animal of claim 1 wherein the non-human transgenic animal is selected from the group consisting of rats, mice, cows, pigs, rabbits, guinea pigs, sheep, hamsters, and goats.
 - 3. A non-human transgenic animal of claim 2 wherein the non-human transgenic animal is further selected from the group consisting of rats and mice.
- 4. A non-human transgenic animal of claim 1 wherein said transgene is operatively linked to a heterologous promoter which effects expression of a primate bradykinin B₁ receptor gene.
- 5. A non-human transgenic animal of claim 4 wherein the non-human transgenic animal is selected from the group consisting of rats, mice, cows, pigs, rabbits, guinea pigs, sheep, hamsters, and goats.
 - 6. A non-human transgenic animal of claim 5 wherein the non-human transgenic animal is further selected from the group consisting of rats and mice.
 - 7. A non-human transgenic animal of claim 4 wherein said heterologous promoter is the rat neuronal specific enolase (NSE) promoter or the cytomegaolvirus (CMV) promoter.
- 30 8. A transgenic rat having incorporated into its genome at least one copy of a transgene encoding a primate bradykinin B₁ receptor gene, or a functional equivalent thereof, such that the transgenic animal demonstrates a humanized B₁ bradykinin receptor occupancy or binding profile.

9. A transgenic rat of claim 8 wherein said transgene is expressed from a heterologous promoter.

- 5 10. A non-human transgenic animal of claim 9 wherein said heterologous promoter is the rat neuronal specific enolase (NSE) promoter or the cytomegaolvirus (CMV) promoter.
- A transgenic rat of claim 10 wherein the transgene is a discistronic
 transgene which further comprises a reporter gene expressed at detectable levels within the transgenic rat.
 - 12. A transgenic rat of claim 11 wherein said transgene is expressed from a heterologous promoter.

13. A non-human transgenic animal of claim 12 wherein said heterologous promoter is the rat neuron specific enolase (NSE) promoter or the cytomegaolvirus (CMV) promoter.

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- 20 14. A transgenic rat having incorporated into its genome at least one copy of a transgene encoding a human bradykinin B₁ receptor gene, or a functional equivalent thereof, such that the transgenic animal demonstrates a humanized B₁ bradykinin receptor occupancy or binding profile.
- 25 15. A transgenic rat of claim 14 wherein said transgene is expressed from a heterologous promoter.
 - 16. A transgenic rat of claim 15 wherein said heterologous promoter is the rat neuronal specific enolase (NSE) promoter or the cytomegaolvirus (CMV) promoter.
 - 17. A transgenic rat of claim 16 wherein the transgene is NSE_CMV intronA_hB1 cds_BGH poly A.

18. A transgenic rat of claim 16 wherein the transgene is CMV_CMVintronA_hB1cds_IRES2_LacZ_BGH polyA.

5 19. A transgenic rat of claim 16 wherein the transgene is Thy1_hB1cds_IRES2_LacZ_BGHpolyA.

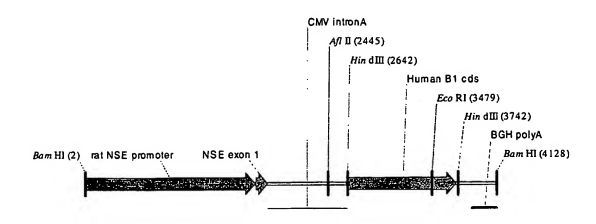
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- 20. An ex vivo method of determining receptor occupancy of a primate bradykinin receptor protein and a test compound, which comprises:
- a) administering the test compound to a non-human transgenic animal, wherein the transgenic animal contains at least one stably integrated copy of a non-native primate B₁ bradykinin gene which expresses levels of non-native primate B₁ bradykinin receptor at levels substantially greater than expression of any endogenous, native B1 bradykinin gene within a target transgenic animal tissue;
 - b) removing a tissue sample from the transgenic animal of step a);
 - c) determining total binding, specific binding and non-specific binding of the test compound to the primate B₁ bradykinin receptor by analysis of the tissue sample of step c);
- d) calculating receptor occupancy of the test compound within the transgenic animal of step a).
 - 21. A method of claim 20 wherein the total binding, specific binding and non-specific binding determination of step c) is accomplished by autoradiography or a tissue homogenate-based binding assay.
 - 22. The method of claim 21 wherein the non-native primate B₁ bradykinin gene is a human B₁ bradykinin gene.
- 23. The method of claim 22 wherein the non-native primate B₁ bradykinin gene is operatively linked to a heterologous promoter.
 - 24. The method of claim 23 wherein said heterologous promoter is the rat neuronal specific enolase (NSE) promoter or the cytomegaolvirus (CMV) promoter.

25. The method of claim 21 wherein the non-human transgenic animal is selected from the group consisting of rats and mice.

- 5 26. The method of claim 25 wherein the non-human transgenic animal is a transgenic rat.
- 27. The method of claim 21 wherein the non-native primate B₁ bradykinin gene is operatively linked to a heterologous promoter and further comprises a
 non-human transgenic animal selected from the group consisting of rats and mice.
 - 28. The method of claim 27 wherein the non-human transgenic animal is a transgenic rat.



NSE hB1 transgene 4132 tp

FIGURE 1

>NSE_CMV intronA_hB1 cds_BGH poly A sequence

GGATCCTGAG	CTCCTCCTCT	GCTCGCCCAA	TCCTTCCAAC	CCCCTATGGT	GGTATGGCTG
	TGTCTGCTCC				
GATGAGGCCT	AGCTTTTGCT	GCTCCAAAGT	TTTAAAAGAA	CACATTGCAC	GGCATTTAGG
	GGTGGAGGAG				
ATCACTGCTT	CCAGGGCCCA	GAGTGGCTTC	CAGGAAGTAT	TCTTACAAAG	GAAGCCCGAT
	CACTCAGAGC				
AGTAACATGA	TCAGTGACCT	GGGGGAGCTG	GCCAAACTGC	GGGACCTGCC	CAAGCTGAGG
GCCTTGGTGC	TGCTGGACAA	CCCCTGTGCC	GATGAGACTG	ACTACCGCCA	GGAGGCCCTG
GTGCAGATGG	CACACCTAGA	GCGCCTAGAC	AAAGAGTACT	ATGAGGACGA	GGACCGGGCA
GAAGCTGAGG	AGATCCGACA	GAGGCTGAAG	GAGGAACAGG	AGCAAGAACT	CGACCCGGAC
	AACCGTACCT				
AAGGTGATGG	CAGGAAGGCA	GTCCCCGGAG	GTCAAAGGCT	GGGCACGCGG	GAGGAGAGGC
CAGAGTCAGA	GGCTGCGGGT	ATCTCAGATA	TGAAGGAAAG	ATGAGAGAGG	CTCAGGAAGA
GGTAAGAAAA	GACACAAGAG	ACCAGAGAAG	GGAGAAGAAT	TAGAGAGGGA	GGCAGAGGAC
CGCTGTCTCT	ACAGACATAG	CTGGTAGAGA	CTGGGAGGAA	GGGATGAACC	CTGAGCGCAT
GAAGGGAAGG	AGGTGGCTGG	TGGTATATGG	AGGATGTAGC	TGGGCCAGGG	AAAAGATCCT
GCACTAAAAA	TCTGAAGCTA	AAAATAACAG	GACACGGGGT	GGAGAGGCGA	AAGGAGGCA
GAGTGAGGCA	GAGAGACTGA	GAGGCCTGGG	GATGTGGGCA	TTCCGGTAGG	GCACACAGTT
CACTTGTCTT	CTCTTTTTCC	AGGAGGCCAA	AGATGCTGAC	GTCAAGAACT	CATAATACCC
CAGTGGGGAC	CACCGCATTC	ATAGCCCTGT	TACAAGAAGT	GGGAGATGTT	CCTTTTTGTC
CCAGACTGGA	AATCCGTTAC	ATCCCGAGGC	TCAGGTTCTG	TGGTGGTCAT	CTCTGTGTGG
CTTGTTCTGT	GGGCCTACCT	AAAGTCCTAA	GCACAGCTCT	CAAGCAGATC	CGAGGCGACT
AAGATGCTAG	TAGGGGTTGT	CTGGAGAGAA	GAGCCGAGGA	GGTGGGCTGT	GATGGATCAG
	AAATAAAAAG	•••		GAGTTCGTGA	
TGGGCTTCTC	CATCTGTCTG	GGTTAGTACC	TGCCACTATA	CTGGAATAAG	GGGACGCCTG
CTTCCCTCGA	GTTGGCTGGA	CAAGGTTATG	AGCATCCGTG	TACTTATGGG	GTTGCCAGCT
		CCTTCCCCCA		TCCCCACCAC	
CGTACGTGCG	TCTCCGCCTG	CAGCTCTTGA			
	ATAGGCGCCG			${\tt CGCGCTGGGA}$	GCCGCAGCCG
	TGCTCTCTCT			GCCACCGCCA	
	CGAGGTAAGT			GCACACCCCT	TTGGCTCTTA
TGCATGCTAT	ACTGTTTTTG	GCTTGGGGCC	TATACACCCC	CGCTTCCTTA	
GATGGTATAG	CTTAGCCTAT	AGGTGTGGGT	TATTGACCAT	TATTGACCAC	TCCCCTATTG
GTGACGATAC		TAATCCATAA		TGCCACAACT	
GCTATATGCC	AATACTCTGT	CCTTCAGAGA	CTGACACGGA	CTCTGTATTT	TTACAGGATG
GGGTCCCATT	TATTATTTAC	AAATTCACAT	ATACAACAAC	GCCGTCCCCC	GTGCCCGCAG
	ACATAGCGTG			GGTACGTGTT	••
	GGTAGCGGCG				
CTCATGGTCG	CTCGGCAGCT	CCTTGCTCCT			
AATGCCCACC		TGCCGCACAA		• • • • • • • • • • • • • • • • • • • •	TGTCTGAAAA
	GATTGGGCTC				
AGAAGATGCA	GGCAGCTGAG	TTGTTGTATT	CTGATAAGAG	TCAGAGGTAA	CTCCCGTTGC

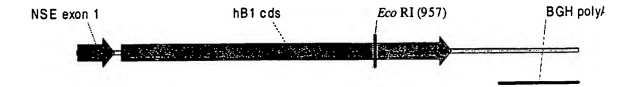
FIGURE 2A

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GGTGCTGTTA ACGGTGGAGG GCAGTGTAGT CTGAGCAGTA CTCGTTGCTG	CCGCGCGCGC
AACCAGAGCC AGCTCTTCCC TCAAAATGCT ACGGCCTGTG ACAATGCTCC	
GACCTGCTGC ACAGAGTGCT GCCGACATTT ATCATCTCCA TCTGTTTCTT	CGGCCTCCTA
GGGAACCTTT TTGTCCTGTT GGTCTTCCTC CTGCCCCGGC GGCAACTGAA	CGTGGCAGAA
ATCTACCTGG CCAACCTGGC AGCCTCTGAT CTGGTGTTTG TCTTGGGCTT	GCCCTTCTGG
GCAGAGAATA TCTGGAACCA GTTTAACTGG CCTTTCGGAG CCCTCCTCTG	CCGTGTCATC
AACGGGGTCA TCAAGGCCAA TTTGTTCATC AGCATCTTCC TGGTGGTGGC	CATCAGCCAG
GACCGCTACC GCGTGCTGGT GCACCCTATG GCCAGCCGGA GGCAGCAGCG	GCGGAGGCAG
GCCCGGGTCA CCTGCGTGCT CATCTGGGTT GTGGGGGGCC TCTTGAGCAT	CCCCACATTC
CTGCTGCGAT CCATCCAAGC CGTCCCAGAT CTGAACATCA CCGCCTGCAT	CCTGCTCCTC
CCCCATGAGG CCTGGCACTT TGCAAGGATT GTGGAGTTAA ATATTCTGGG	TTTCCTCCTA
CCACTGGCTG CGATCGTCTT CTTCAACTAC CACATCCTGG CCTCCCTGCG	AACGCGGGAG
GAGGTCAGCA GGACAAGGTG CGGGGGCCGC AAGGATAGCA AGACCACAGC	GCTGATCCTC
ACGCTCGTGG TTGCCTTCCT GGTCTGCTGG GCCCCTTACC ACTTCTTTGC	CTTCCTGGAA
TTCTTATTCC AGGTGCAAGC AGTCCGAGGC TGCTTTTGGG AGGACTTCAT	TGACCTGGGC
CTGCAATTGG CCAACTTCTT TGCCTTCACT AACAGCTCCC TGAATCCAGT	AATTTATGTC
TTTGTGGGCC GGCTCTTCAG GACCAAGGTC TGGGAACTTT ATAAACAATG	CACCCCTAAA
AGTCTTGCTC CAATATCTTC ATCCCATAGG AAAGAAATCT TCCAACTTTT	CTGGCGGAAT
TAAAACAGCA TTGAACCAAG AAGCTTGGCT TTCTTATCAA TTCTTTGTGA	CATAATAAAT
GCTATTGTGA TAGGCTAAAT GATTACTCCC GTAGATTGGG GGGTACCTAA	TCCCTGGACT
TGATGAGCGG CCTCGAGCAT GCATCTAGAG GGCCCTATTC TATAGTGTCA	CCTAAATGCT
AGAGCTCGCT GATCAGCCTC GACTGTGCCT TCTAGTTGCC AGCCATCTGT	TGTTTGCCCC
TCCCCCGTGC CTTCCTTGAC CCTGGAAGGT GCCACTCCCA CTGTCCTTTC	CTAATAAAAT
GAGGAAATTG CATCGCATTG TCTGAGTAGG TGTCATTCTA TTCTGGGGGG	TGGGGTGGGG
CAGGACAGCA AGGGGGAGGA TTGGGAAGAC AATAGCAGGC ATGCTGGGAT	CC (SEQ ID
NO:1)	

FIGURE 2B

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NSE hB1 transcript

FIGURE 3A

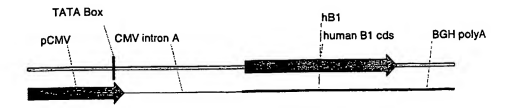
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GCTTGGTACC	GCCGCCACCA	TGGCATCATC	CTGGCCCCCT	CTAGAGCTCC	AATCCTCCAA
CCAGAGCCAG	CTCTTCCCTC	AAAATGCTAC	GGCCTGTGAC	AATGCTCCAG	AAGCCTGGGA
CCTGCTGCAC	AGAGTGCTGC	CGACATTTAT	CATCTCCATC	TGTTTCTTCG	GCCTCCTAGG
GAACCTTTTT	GTCCTGTTGG	TCTTCCTCCT	GCCCCGGCGG	CAACTGAACG	TGGCAGAAAT
CTACCTGGCC	AACCTGGCAG	CCTCTGATCT	GGTGTTTGTC	TTGGGCTTGC	CCTTCTGGGC
AGAGAATATC	TGGAACCAGT	TTAACTGGCC	TTTCGGAGCC	CTCCTCTGCC	GTGTCATCAA
CGGGGTCATC	AAGGCCAATT	TGTTCATCAG	CATCTTCCTG	GTGGTGGCCA	TCAGCCAGGA
CCGCTACCGC	GTGCTGGTGC	ACCCTATGGC	CAGCCGGAGG	CAGCAGCGGC	GGAGGCAGGC
CCGGGTCACC	TGCGTGCTCA	TCTGGGTTGT	GGGGGGCCTC	TTGAGCATCC	CCACATTCCT
GCTGCGATCC	ATCCAAGCCG	TCCCAGATCT	GAACATCACC	GCCTGCATCC	TGCTCCTCCC
CCATGAGGCC	TGGCACTTTG	CAAGGATTGT	GGAGTTAAAT	ATTCTGGGTT	TCCTCCTACC
ACTGGCTGCG	ATCGTCTTCT	TCAACTACCA	CATCCTGGCC	TCCCTGCGAA	CGCGGGAGGA
GGTCAGCAGG	ACAAGGTGCG	GGGGCCGCAA	GGATAGCAAG	ACCACAGCGC	TGATCCTCAC
GCTCGTGGTT	GCCTTCCTGG	TCTGCTGGGC	CCCTTACCAC	TTCTTTGCCT	TCCTGGAATT
CTTATTCCAG	GTGCAAGCAG	TCCGAGGCTG	CTTTTGGGAG	GACTTCATTG	ACCTGGGCCT
GCAATTGGCC	AACTTCTTTG	CCTTCACTAA	CAGCTCCCTG	AATCCAGTAA	TTTATGTCTT
TGTGGGCCGG	CTCTTCAGGA	CCAAGGTCTG	GGAACTTTAT	AAACAATGCA	CCCCTAAAAG
TCTTGCTCCA	ATATCTTCAT	CCCATAGGAA	AGAAATCTTC	CAACTTTTCT	GGCGGAATTA
AAACAGCATT	GAACCAAGAA	GCTTGGCTTT	CTTATCAATT	CTTTGTGACA	TAATAAATGC
TATTGTGATA	GGCTAAATGA	TTACTCCCGT	AGATTGGGGG	GTACCTAATC	CCTGGACTTG
ATGACGCTCG	AGCATGCATC	TAGAGGGCCC	TATTCTATAG	TGTCACCTAA	ATGCTAGAGC
TCGCTGATCA	GCCTCGACTG	TGCCTTCTAG	TTGCCAGCCA	TCTGTTGTTT	GCCCCTCCCC
CGTGCCTTCC	TTGACCCTGG	AAGGTGCCAC	TCCCACTGTC	CTTTCCTAAT	AAAATGAGGA
AATTGCATCG	CATTGTCTGA	GTAGGTGTCA	TTCTATTCTG	GGGGGTGGGG	TGGGGCAGGA
CAGCAAGGGG	GAGGATTGGG	AAGACAATAG	CAGGCATGCT	GGGGA (SEQ	ID NO:2)

FIGURE 3B

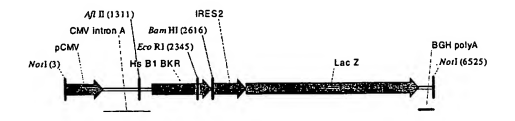
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CMV Promoter - CMV Intron A-human B1 cds-BGH
2986 bp

FIGURE 4

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pCMV B1 IZ insert 6532 bp

FIGURE 5

8/16 >promoterCMV_CMVintronA_hB1cds_IRES2_LacZ_BGH polyA

	GTACGGGCCA				
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TACGGTAAAT	GGCCCGCCTG	GCTGACCGCC	CAACGACCCC	CGCCCATTGA	CGTCAATAAT
GACGTATGTT	CCCATAGTAA	CGCCAATAGG	GACTTTCCAT	TGACGTCAAT	GGGTGGAGTA
TTTACGGTAA	ACTGCCCACT	TGGCAGTACA	TCAAGTGTAT	CATATGCCAA	GTACGCCCCC
TATTGACGTC	AATGACGGTA	AATGGCCCGC	CTGGCATTAT	GCCCAGTACA	TGACCTTATG
GGACTTTCCT	ACTTGGCAGT	ACATCTACGT	ATTAGTCATC	GCTATTACCA	TGGTGATGCG
GTTTTGGCAG	TACATCAATG	GGCGTGGATA	GCGGTTTGAC	TCACGGGGAT	TTCCAAGTCT
CCACCCCATT	GACGTCAATG	GGAGTTTGTT	TTGGCACCAA	AATCAACGGG	ACTTTCCAAA
ATGTCGTAAC	AACTCCGCCC	CATTGACGCA	AATGGGCGGT	AGGCGTGTAC	GGTGGGAGGT
CTATATAAGC	AGAGCTCTCT	GGCTAACTAG	AGAACCCACT	GCTTACTGGC	TTATCGAAAT
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	CTATACTGTT				
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	ATACTTTCCA				
	TGCCAATACT				
	CATTTATTAT				
	TTAAACATAG				
	CTCCGGTAGC				
	GTCGCTCGGC				
	CACCACCACC				
	TGGAGATTGG				
	TGCAGGCAGC				
	GTTAACGGTG				
	ACATAATAGC				
	GGTACCGCCG				
	AGCCAGCTCT				
	CTGCACAGAG				
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AGAAATCTAC			TGATCTGGTG		
	AATATCTGGA				_
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•	TACCGCGTGC				
	GTCACCTGCG				
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TTCCACCATA	TTGCCGTCTT	TTGGCAATGT	GAGGGCCCGG	AAACCTGGCC	CTGTCTTCTT
GACGAGCATT	CCTAGGGGTC	TTTCCCCTCT	CGCCAAAGGA	ATGCAAGGTC	TGTTGAATGT
CGTGAAGGAA					
	CGGAACCCCC				
ATAAGATACA					
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GGTACCCCAT					
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GGGAAAACCC					
GGCGTAATAG					

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		CAGGACAGTC			
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GCCGCGCTGT	ACTGGAGGCT	GAAGTTCAGA	TGTGCGGCGA	GTTGCGTGAC	TACCTACGGG
TAACAGTTTC	TTTATGGCAG	GGTGAAACGC	AGGTCGCCAG	CGGCACCGCG	CCTTTCGGCG
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		GGTAACAGTC			TGGCAGGCGT
GCGAATACGC	CCACGCGATG	CAGGGCGGCT	TIGGCGGIII	COCIAMIAC	
		AACCCGTGGT			
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		ATGAACGGTC			
		CAGCAGCAGT			GGGCAAACCA
		CTGTTCCGTC			CACTGGATGG
		CTGGCAAGCG			
		CCTGAACTAC			
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		CAGTTCACCC			
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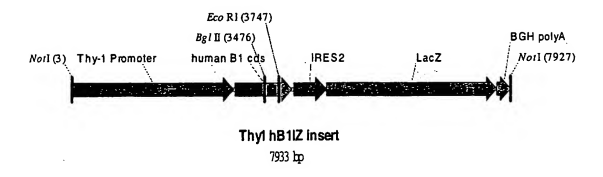


FIGURE 7

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CAGCAAGCCA	CTTCGGACAC	CCAAATGGAA	CACCTCCAGT	CAGCCCTCGC	CGACCACCCC
ACCCCCTCCA	TCCTTTTCCC	TCAGCCTCCG	ATTGGCTGAA	TCTAGAGTCC	CTCCCTGCTC
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		GCATCAGGGT			TCAGGGAAAG
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ACATCCTGAG	ATACATTGGG	AGAGGAGATA	CAGTTTCAAT	AAAATAATAG	GTTGGGTGGA
		CCACTCAGGA		GCTTCGTGAG	TTTGAGGCCA
	CATAGTGAAA	CCCTGTCAGT	AAATAAGTAA	GCAAGTATTT	GAGTATCTAC
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		ACACCCCTAA		CAGACCAATT	AGACAATAAA
	AGCTTACCAA		TGTATTTTCA		TGTCTGTGTA
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CCTCTGCCTC	TCTGCCTCTC		CCTCTCTCTC	TGCCTCTCTC	TGCCTCTCTC
TGCCCCTCTC	TGCCCCTCTC	TGCCCCTCTC	TGCCCCTCTC	TGCCGCCCTC	TGCCTTCTGC
CCTCTGCCCT	CTGGCCTCTG	GCCTCTGCCC	TCTGCCCTCT	GGCCTCTGGC	CTCTGCCTCT
••			GCTCTGTAGG	TCTTAAGTTC	CAGAAGAAAG
GCCTCTTGAG	TGCTGGAATC				CCAGGACATA
TAATGAAGTC		GAGGTGCTCA		AGACACACAC	
GGCTCCCACT	TCCTTGGCTT		GCAAAGGACC	TTAGGCAGTG	TCACTCCCTA
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GCCCCGGCGG	CAACTGAACG	TGGCAGAAAT	CTACCTGGCC	AACCTGGCCG	CCTCTGATCT
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				ATCCAAGCCG	
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	AATCCAGTAA			CTCTTCAGGA	CCAAGGTCTG
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	CCCTCCCCC			AGCCGCTTGG	
		TATTTTCCAC		TCTTTTGGCA	
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	TCTGTAGCGA			CCCCCACCTG	
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	TCGAGGCGTT			TCTGCATGGT	
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GGGACTGGGT	GGATCAGTCG	CTGATTAAAT	ATGATGAAAA	CGGCAACCCG	TGGTCGGCTT
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""COCCOTON					

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ATAACGAGCT	CCTGCACTGG	ATGGTGGCGC	TGGATGGTAA	GCCGCTGGCA	AGCGGTGAAG
TGCCTCTGGA	TGTCGCTCCA	CAAGGTAAAC	AGTTGATTGA	ACTGCCTGAA	CTACCGCAGC
CGGAGAGCGC	CGGGCAACTC	TGGCTCACAG	TACGCGTAGT	GCAACCGAAC	GCGACCGCAT
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CGCTGGATAA	CGACATTGGC	GTAAGTGAAG	CGACCCGCAT	TGACCCTAAC	GCCTGGGTCG
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FIGURE 8C

14/16 Rat 1810

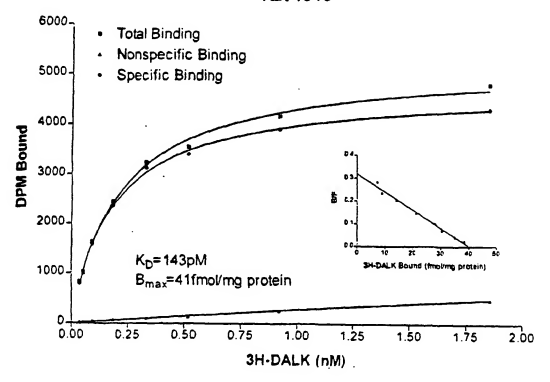
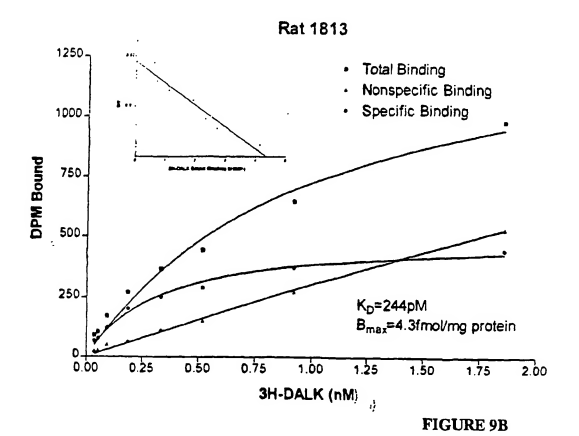


FIGURE 9A



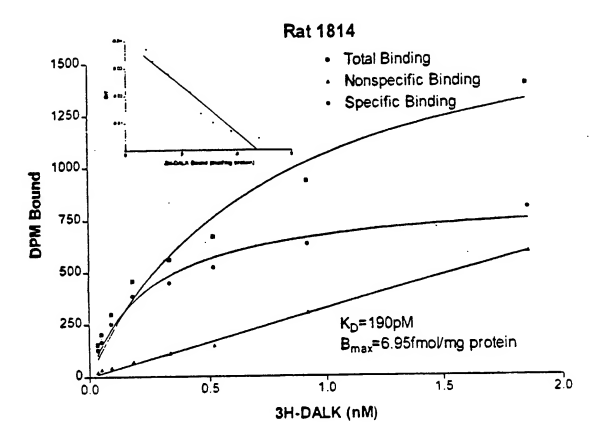


FIGURE 9C

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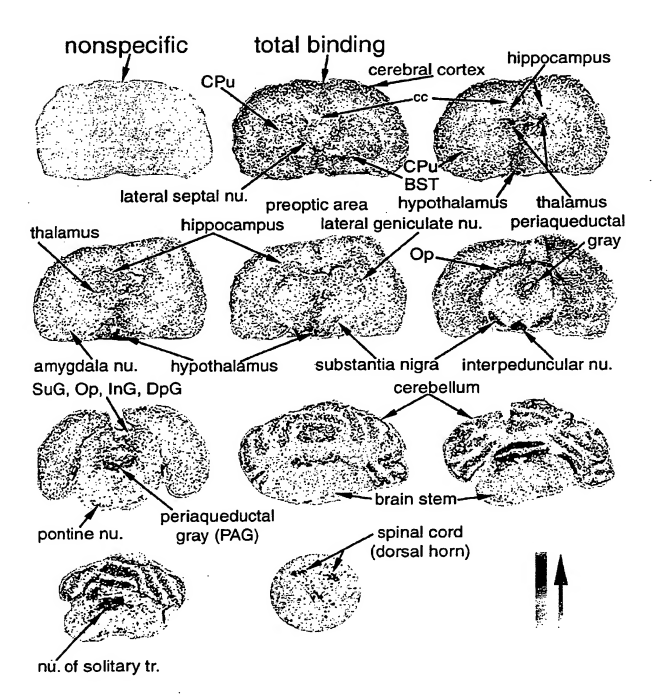


FIGURE 10

SEOUENCE LISTING

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